This Listing of Claims will replace all prior versions, including listings, of claims in the

application.

Listing of Claims

Claim 1 (currently amended): A method to enable the assessment of the growth rate and

death rate of a micro-organism within a chosen time period in an environment of interest by

introducing into said micro-organism at least two reporter genes, which method is characterised in

that

a) said reporter genes code for luminescent and/or fluorescent products and within said time

period and environment at least two said products of the following are produced:

i) an essentially a stable product produced in a step (a), within the environment of

interest, essentially known proportion to the total amount of cells of said micro-organism that are

or have been alive within said chosen time period,

ii) a product present in said environment of interest in an essentially known

proportion to the amount of cells alive at any moment within said chosen time period, and

iii) an essentially a stable product produced in a step (a), within the environment of

interest, essentially known proportion to the total amount of cells of said micro-organism that have

died within said chosen time period,

and said products can be measured through their luminescence and/or fluorescence;

b) said micro-organism is incubated within the environment of interest and said

luminescence and/or fluorescence is detected after said chosen time period, and

c) the growth and death rate of the said micro-organism is assessed based on at least two of

the following:

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i) the known proportion of luminescence or fluorescence to the amount of cells alive

after any said chosen time period,

ii) the known proportion of luminescence or fluorescence to the total amount of cells

that are or have been alive within any said chosen time period, and

iii) the known proportion of luminescence or fluorescence to the total amount of cells

that have died within any said chosen time period.

Claim 2 (original): The method according to claim 1 characterised in that said micro-

organism is a gram negative bacteria, e.g. Escherichia coli.

Claim 3 (previously presented): The method according to claim 1 characterised in that

a) one reporter gene coding for a luminescent product is luciferase, which is used for the

determination of amount of cells alive at any moment within said chosen time period, and

b) another reporter gene coding for a fluorescent product is green fluorescent protein (GFP),

which is used for the determination of total amount of cells of said micro-organism that are or have

been alive within said chosen time period.

Claim 4 (previously presented): The method according to claim 1 characterised in that said

reporter genes are introduced into said micro-organism in a plasmid.

Claim 5 (previously presented): The method according to claim 3 characterised in that said

plasmid is pGFP+luc* (SEQ ID NO: 1).

Claim 6 (previously presented): The method according to claim 2 characterised in that

a) one reporter gene coding for a luminescent product is luciferase, which is used for the

determination of amount of cells alive at any moment within said chosen time period, and

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b) another reporter gene coding for a fluorescent product is green fluorescent protein (GFP), which is used for the determination of total amount of cells of said micro-organism that are or have been alive within said chosen time period.

Claim 7 (previously presented): The method according to claim 2 characterised in that said reporter genes are introduced into said micro-organism in a plasmid.

Claim 8 (previously presented): The method according to claim 4 characterised in that said plasmid is pGFP+luc* (SEQ ID NO: 1).

Claim 9 (previously presented): The method according to claim 6 characterised in that said plasmid is pGFP+luc* (SEQ ID NO: 1).

Claim 10 (previously presented): The method according to claim 7 characterised in that said plasmid is pGFP+luc* (SEQ ID NO: 1).